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TITLE OF INVENTION: PROCESS FOR BASE EXCHANGE OF PHOSPHOLIPIDS WHAT IS CLAIMED IS:

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1. A process for the base exchange of a phospholipid with an enzyme. which comprises reacting phosphatidylcholine as the phospholipid with an alcohol selected from the group consisting of sering ethanolamine, Nmethylethanolamine, N,N-dimethylethanolamine, glycerol and monosaccharides in the presence of phospholipase D of Streptomyces origin as the enzyme so as to accomplish the base exchange reaction. المدران روق مدرو

DETAILED EXPLANATION OF INVENTION:

(Related Technical Field)

The present invention relates to a process for the base exchange of phospholipids using an enzyme. In particular, it relates to a process for the base exchange comprising treating phosphatidylcholing with phospholipase D.

(Problem Underlining Invention)

As to the base exchange of phospholipids with enzymes, there is known a process wherein phosphatidylcholine is treated with phospholipase D for the base exchange so as to obtain a phospholipid comprising a desired base (S.F. Yang et al., J.Biol.Chem., 242 (3), 477-484 (1967); R.M.C. Dawson, Biochem. J.. 102, 205-210 (1967)].

In said process, the base exchange is carried out mainly by the use of phospholipase D of cabbage origin, but the conversion is less than 13 %. Also, the alcohol usable for the base exchange is limited to a primary alcohol of not more than 5 carbon atoms. Especially, the conversion for nitrogen-containing alcohols is very low and 12 % at the highest. In case of monosaccharides, no base exchange is observed. This invention is directed to improvement in the above respects, i.e. expansion of the range of usable alcohols and attainment of the base exchange of phospholipids with high yields of the desired products.

(Solution for Technical Problem)

The present invention is a process for the base exchange of phospholipids with an enzyme which comprises reacting phosphatidylcholine as the substrate with an alcohol selected from the group consisting of serine, ethanolamine, N-methylethanolamine, N,N-dimethylethanolamine, glycerol and monosaccharides in the presence of phospholipase D of Streptomyces origin.

The phosphatidylcholine usable in this invention may be either a natural product obtained by extraction and purification or a synthetic product.

The phospholipase D of Streptomyces origin asable in this invention is the one obtainable from phospholipase D-producing microorganisms such as Streptomyces chromofoscus or the like and may be commercially available on

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the market.

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The reaction can be carried out by contacting phosphatidylcholine with an alcohol in the presence of phospholipase D of Streptomyces origin. As the alcohol, there may be used the one chosen from nitrogen-containing alcohols such as serine, ethanolamine, N-methylethanolamine and N,N-dimethylethanolamine as well as polyols such as glycerol and monosaccharides. As the monosaccharides, aldoses and ketoses may be used, and their examples are pentoses such as ribose and arabinose, hexoses such as glucose and fructose, etc.

The solvent to be used for the reaction is water alone or a mixture of water and an organic solvent. Examples of the organic solvent are aliphatic hydrocarbons (e.g. n-heptane, n-hexane, petroleum ether), alicyclic hydrocarbons (e.g. cyclopentane, cyclohexane), ethers (e.g. diethyl ether, tetrahydrofuran), esters (e.g. methyl acetate, ethyl acetate), halogenated hydropearbons (e.g. carbon tetrachloride, chloroform), etc. When a mixture of water and an organic solvent is used, their proportion may be optional; for instance, the weight ratio of water and the organic solvent may be from 1:1 to 0.1:10. For suppression of the side-reaction and obtainment of the objective product in a high yield, it is preferred to keep the water content in the reaction system not more than 10 % by weight.

The molar ratio of phosphatidylcholine and the alcohol may be appropriately chosen depending on the kind of the alcohol; in general, the alcohol may be used in an amount of 5 - 100 mol to 1 mol of phosphatidylcholine.

The amount of phospholipase D of Streptomyees origin to be used may be selected from the range of 100 ~ 500 units per 1g of phosphatidylcholine.

After the reagents are charged as above, the resultant mixture is reacted, for instance, at a temperature of 20 to 60°C while stirring with rotation or supersonic waves for a period of 30 minutes to 5 hours.

(Effect of Invention)

In the present invention, the base exchange reaction of phosphatidylcholine with phospholipase D is carried out by the use of phospholipase D of Streptomyces origin instead of phospholipase D of cabbage origin, whereby a phospholipid substituted with a desired alcohol can be obtained in a high yield. Also, it is possible to accomplish the base exchange with a monosaccharide such as glucose, which could not be accomplished by the use of phospholipase D of cabbage origin. Accordingly, the present invention makes the scope of the alcohol as exchangeable expanded and the conversion of the phospholipid improved.

The present invention will be explained more in details by way of practical embodiments.

Example 1

One hundred ut of a suspension prepared by suspending 40 mg of dipalmitoylphosphatidylcholine in 1 ml of water, 0.5 M acetate buffer (pH 5) (50 ut), ethanolamine (adjusted to pH 5 with 0.5 N HCi) (50 mg), diethyl ether (1 ml),

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an aqueous solution of phospholipase D of Streptomyces origin ("Phospholipase DP" manufactured by Toyo Jozo)(50 units/ml)(10 ul) were mixed at 37°C while stirring at 500 rpm for 1 hour, and after completion of the reaction, the phosholipid was extracted with chloroform.

The extract was subjected to analysis with thin layer chromatography using a developing solvent of chloroform; acctone; methanol; acctic acid; water = 50; 20; 10: 15: 5 and the Dittmer reagent as a color developing agent, and the composition of the product was determined by densitometry.

As the result, it was revealed that the product comprised 90 % of phosphatidylethanolamine and 10 % of phosphatidic and.

Example 2

The reaction was carried out in the same manner as in Example 1 but using scrine (100 mg) in place of ethanolamine. After completion of the reaction, extraction and analysis were carried out in the same manner as in Example 1. As the result, the product comprised 70 % of phosphatidylsenine. 20 % of phosphatidic acid and 10 % of phosphatidylcholine.

Example 3

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The reaction was carried out in the same marner as in Example 1 but using glucose (150 mg) in place of ethanolamine. After completion of the reaction, extraction and analysis were carried out in the same manner as in Example 1. As the result, the product comprised 63 % of phosphatidylglucose, 21 % of phosphatidic acid and 16 % of phosphatidylcholine.

Example 4

The reaction was carried out in the same manner as in Example 1 but using glycerol (70 mg) in place of ethanolamine. After completion of the reaction, extraction and analysis were carried out in the same manner as in Example 1. As the result, the product comprised 81 % of phosphatidylglycerol, 11 % of phosphatidic acid and 8 % of phosphatidylcholine.

Example 5

The reaction was carried out in the same manner as in Example 1 but using phospholipase D of Streptomyces chromofoscus (manufactured by Bohringer Mannheim) in place of phospholipase D of Streptomyces origin ("Phospholipase DP" manufactured by Toyo Jozo). After completion of the reaction, extraction and analysis were carried out in the same manner as in Example 1. As the result, the product comprised 83 % of phosphatidylethanolamine, 10 % of phosphatidic acid and 7 % of phosphatidyletholine.

Example 6

The reaction was carried out in the same manner as in Example 1 but using glucose (150 mg) in place of ethanolamine. After completion of the reaction, extraction and analysis were carried out in the same manner as in Example 1. As the result, the product comprised 51 % of phosphatidylglucose. 12 % of phosphatidic acid and 37 % of phosphatidylcholine.

Comparative Examples 1 to 4

The reaction was carried out in the same manner as in Examples 1 to 4 but using phospholipase D of cabbage origin in place of phospholipase D of

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Streptomyces origin and adding thereto calcium chloride (10 mg) for activation, the alcohol used being ethanol (Comparative Example 1), serine (Comparative Example 2), glucose (Comparative Example 3) or glycerol (Comparative Example 4).

The analytical results in Examples and Comparative Examples as above are shown in the following table:

	Product	Phosphatidic acid	Phosphatidy) choline
Example 1	90%	10%	0 %
Comparative 1	40%	55%	5%
Example 2	70%	20%	10%
Comparative 2	0%	52%	48%
Example 3	63%	21%	16%
Comparative 3	0%	60%	40%
Example 4	81%	11%	8%
Comparative 4	26%	66%	8%
Example 5	83%	. 10%	7%
Example 6	51%	12%	37%

As understood from the above table, the yields of the products in Examples are markedly higher than those in Comparative Examples for all the alcohols.

In Comparative Examples where phospholipase D of cabbage origin is used, the conversion into phosphatidylethanolamine or phosphatidylelycerol as the exchange reaction product in case of using ethanolamine (Comparative Example 1) or glycerol (Comparative Example 4) as the alcohol is lower than that in Example 1 or 4. It is especially noted that when the alcohol is serine (Comparative Example 2) or glucose (Comparativ3e Example 3), phosphatidylserine or phosphatidylglucose as the product was not obtained.

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発明の数 1 (全3頁)

❷発明の名称 リン語質の収益交換反応法

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 $\Theta \Delta$ **昭63--3679**D

顧 昭61(1986) 8月1日

@昭63(1988) 2月17日

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の特許青水の範囲

17 : 酵素を利用してリン脂質の塩基交換反応を行 うに殴し、基質としてホスファチジルコリンを用 い、セリン、エタノールアミン、Nーメチルエタ ン、グリセロールおよび単糖の群から選ばれるア ルコールを、ストレブトマイセス属由来のホスホ リバーゼDの存在下に反応せしめ、塩基交換を進 行させることを特徴とするリン脳質の塩基交換反 定准。

発明の詳細な歴明

(廃棄上の利用分野)

本発明は酵素を利用したリン脳質の塩素交換反 応法に関し、特にホスフアチジルコリンにホスホ リパーゼDを作用させる塩基交換反応法に関す 15 反応を行うに廃し、基質としてホスファチジルコ

(従来の技術と発明が解決しようとする問題点) 酵素を利用したリン脂質の塩基交換反応におい て、ホスフアチジルコリンにホスホリバーゼDを 作用させ、塩基交換反応法により目的とする塩基 20 のホスポリバーゼロの存在下に反応せしめ、塩基 を持つリン酸質を製造する技術は公知である。 (S.F.Yang, et al., J.Biol.Chem.242, (3) 477 - 484(1967)), [R.M.C. Dawson, Biochem, J.

102, 205~210(1967)]

これらの批領では、主としてキャベッ由来ホス ホリバーゼDを用いて塩塞交換反応を行っている が、その変換率は13%以下であった。また交換反 ノールアミン、N, Nージメチルエタノールアミ 5 応に使用できるアルコールは、炭素数 5以下の一 歴アルコールに限られていた。特に、含窒素アル コールに関しては変換率が非常に低く、高いもの でも12%であつた。また、単階については交換反 応が認められなかつた。本発明は、これらの点を 10 改善し、使用できるアルコールの範囲を広げ、し かも高収率で目的物が得られるリン脳質の塩基交 換反応法を提供することを目的とする。

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(問題点を展決するための手段)

本発明は、酵素を利用してリン脂質の塩基交換 リンを用い、セリン、エタノールアミン、Nーメ チルエタノールアミン、N, Nージメチルエタノ ールアミン、グリセロールおよび単特の群から選 ばれるアルコールを、ストレブトマイセス医由来 交換を進行させることを特徴とするリン脂質の塩 基交換反応法である。

本発明で使用するホスフアチジルコリンは、天

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ミリモル加え、その他の反応条件、操作は実施例 1と全く同様に行った。

分析権果は、ホスフアチジルグルコース四名。 ホスフアチジン酸21%。ホスフアチジルコリン16 光であった。

爽遊例 4

エタノールアミンのかわりにグリセロールを70 ミリモル加え、その他の反応条件、操作は実施例 1と全く同様に行った。

分析結果は、ホスフアチジルグリセロール81 10 %、ホスフアチジン酸11%、ホスフアチジルコリ ン8%であった。

突旅例 5

ストレプトマイセス協由来のホスホリバーゼD (東洋顔造器製のホスホリバーゼDP) のかわりに 15 ストレブトマイセス・クロモポスカス由来のホス ホリパーゼD(ベーリンガーマンハイム社製) に 変えた以外は、実施例1と同様の組成、条件で反 応を行った。反応終了後天施例1と同じ操作にて 抽出、分折を行った。

その結果、ホスフアチジルエタノールアミン83 %,ホスフアチジン酸10%,ホスフアチジルコリ ン7%であった。

実施例 6

にグルコースを150ミリモル加土た以外は、同様 の組成、条件で反応を行った。反応終了後実施例 1と同じ操作にて抽出、分析を行った。

その結果、ホスフアナジルグルコース51%。ホ スフアチジン酸12%。ホスフアチジルコリン87% 30 ホスフアチジルセリンおよびホスフアチジルグル であった。

比较例 1~4

ストレプトマイセス医由来のホスホリバーゼD のかわりにキャベツ由来ホスホリバーゼDを用 い、活性化の為に塩化カルシウムを10ミリモルを 35 加えた以外は、添加アルゴールとしてエタノール アミン(比較例1)、セリン(比較例2)、グルコ

ース(比較例3)、グリセロール(比較例4)を 用い、実施例1~4までと同様に行った。 実施例および比較例の分析結果を下表に示す。

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	生成 初	ホスフア チジン酸	ホスフアチ ジルコリン
FEE (F)	90%	10%	0%
比較例1	40%	55%	5%
実施例 2.	70%	20%	10%
比較例 2	0%	52%	48%
美政例3	63%	21%	16%
比較好3	0%	60%	40%
类施例 4	81%	11%	8%
土較例4	26%	55%	8%
突旋例 5	83%	10%	796
実施例 6	51%	12%	37%

表から切らかなように、実施例では、比较例よ 20 りも、すべて各アルコールについて著しく高収率 で目的物が得られる。

これに対して比較例では、キャベツ由来ホスホ リパーゼDを使用したので、添加アルコールがエ タノールアミン(比較例1)およびグリセロール **実施例 5 において、エタノールアミンのかわり 25 (比較例 4) の場合、交換反応による生成物、ホ** スフアチジルエタノールアミンおよびホスファチ ジルグリセロールへの変換率が実施例1および4 に比較して低い。特に、セリン(比較例2)およ びグルコース(比較例3)の場合には、生成物の コースが全く係られなかつた。